(REV. 1-98) TRANSMITTAL LETTER TO THE UNITED STATES 0696-0171P DESIGNATED/ELECTED OFFICE (DO/EO/US) CONCERNING A FILING UNDER 35 U.S.C. 371 INTERNATIONAL APPLICATION NO. INTERNATIONAL FILING DATE PRIORITY PCT/FI98/01004 December 21, 1998 December 23, 1997 TITLE OF INVENTION A SOLUBLE MAGNESIUM DIHALIDE COMPLEX, ITS PREPARATION AND USE APPLICANT(S) FOR DO/EO/US GAROFF, Thomas; LEINONEN, Timo; ALA-HUIKKU, Sirpa Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information: This is a FIRST submission of items concerning a filing under 35 U.S.C. 371. This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U.S.C. 371. This express request to begin national examination procedures (35 U.S.C. 371(f)) at any time rather than delay examination until the expiration of the applicable time limit set in 35 U.S.C. 371(b) and PCT Articles 22 and 39 (1). A proper Demand for International Preliminary Examination was made by the 19th month from the earliest claimed priority date A copy of the International Application as filed (35 U.S.C. 371(c)(2)) is transmitted herewith (required only if not transmitted by the International Bureau). has been transmitted by the International Bureau. is not required, as the application was filed in the United States Receiving Office (RO/US). A translation of the International Application into English (35 U.S.C. 371(c)(3)). Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(2)). are transmitted herewith (required only if not transmitted by the International Bureau). O have been transmitted by the International Bureau. have not been made; however, the time limit for making such amendments has NOT expired. 111 d. A have not been made and will not be made. A translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)). An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)). A translation of the annexes to the International Preliminary Examination Report under PCT Article 36 10. (35 U.S.C. 371(c)(5)). Items 11. to 16. below concern document(s) or information included: An Information Disclosure Statement under 37 CFR 1.97 and 1.98.-PTO-1449/International Search Report (PCT/ISA/210) An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included. A FIRST preliminary amendment. A SECOND or SUBSEQUENT preliminary amendment. A substitute specification. A change of power of attorney and/or address letter. 16. Other items or information: 1.) Preliminary Examination Report (PCT/IPEA/409) 2.) PCT Request (PCT/RO/ 101)

S. APPLICATION NO (af known, see 37 CFR 1

3.) Five (5) sheets of Formal Drawings

INTERNATIONAL APPLICATION NO

ATTORNEY'S DOCKET NUMBER

PCT/F198/01004

0696-0171P

an Iran	771	430	THECOM	Ui	TIU 23	JUN ZUUD
17. The full win lies	ne su mitted:				LCULATIONS	PTO USE ONLY
	FEE (37 CFR 1.492(a)(1)-(5): reliminary examination fee (37)					
	h fee (37 CFR 1.445(a)(2)) pai					
and International Sear	ch Report not prepared by the	EPO or JPO	\$970.00			
	ry examination fee (37 CFR 1 nal Search Report prepared by		\$840.00			
	ary examination fee (37 CFR 1 n fee (37 CFR 1.445(a)(2)) pai		\$690.00			
	ary examination fee (37 CFR 1 atisfy provisions of PCT Artic		\$670.00			
	ry examination fee (37 CFR 1		\$96.00	-		E
	provisions of PCT Article 33(PROPRIATE BASIC		390.00	\$	840.00	
	or furnishing the oath or declar		∑ 30	\$	130.00	
months from the earlies	at claimed priority date (37 CF) NUMBER FILED	R 1.492(e)). NUMBER EXTRA	RATE	_		L
Total Claims	32 - 20 =	12	X \$18.00	\$	216.00	
Independent Claims	1 - 3 =	0	X \$78.00	\$	0.00	
	ENT CLAIM(S) (if applicable)		+ \$260.00	\$	0.00	
		OF ABOVE CALCULA		\$	1186.00	
	g by small entity, if applicable			\$	0.00	
must also be filed (Note	2 37 CFR 1.9, 1.27, 1.28).	CIID	TOTAL -			
Brocessing fee of \$130	00 for furnishing the English t		FOTAL = 20 30	\$	1186.00	
	t claimed priority date (37 CF)		20 <u> </u>	\$	0.00	
Street,		TOTAL NATIONA		\$	1186.00	
	closed assignment (37 CFR 1. ropriate cover sheet (37 CFR 3			\$	0.00	
		TOTAL FEES ENC		\$	1186.00	
					Amount to be:	\$
11. 174				┝	refunded charged	S
a. ⊠ A check in the an	nount of \$ 1186.00 to cover th	ne above fees is enclosed.				
b. ☐ Please charge my		in the amount of \$	to co	ver	the above fees	
	of this sheet is enclosed.					
	er is hereby authorized to char Deposit Account No. <u>02-2448</u>		h may be req	uire	d, or credit any	
	ppropriate time limit under t be filed and granted to rest			et, a	petition to revi	ve (37 CFR
Send all correspondence to: Birch, Stewart, Kola P.O. Box 747	sch & Birch, LLP or Custo	mer No. 2292	SIGNATU	RE	/_w	y
Falls Church, VA 22 (703)205-8000	040-0747				NDREW D.	
			#32,868			
/cw June 23, 2000			REGISTRA	ATION	NUMBER	

A STATE OF THE STA

09/58232**1** 430 Rec'd PCT/PTO 23 JUN **2000**

PATENT 0696-0171P

IN THE U.S. PATENT AND TRADEMARK OFFICE

Applicant:

GAROFF, Thomas et al.

Int'l. Appl. No.:

PCT/FI98/01004

Appl. No.:

new

Group:

UNKNOWN

Filed:

June 23, 2000

Examiner: UNKNOWN

For:

A SOLUBLE MAGNESIUM DIHALIDE COMPLEX, ITS PREPARATION AND USE

PRELIMINARY AMENDMENT

BOX PATENT APPLICATION

Assistant Commissioner for Patents Washington, DC 20231

June 23, 2000

Sir:

The following Preliminary Amendments and Remarks are respectfully submitted in connection with the above-identified application.

AMENDMENTS

IN THE SPECIFICATION:

Please amend the specification as follows:

Before line 1, insert --This application is the national phase under 35 U.S.C. § 371 of PCT International Application No. PCT/FI98/01004 which has an International filing date of December 21, 1998, which designated the United States of America.--

IN THE CLAIMS:

Please amend the claims as follows:

CLAIM 3: Line 1, delete "or 2"

CLAIM 4: Line 1, delete ", 2 or 3"

CLAIM 5: Line 1, change "any preceding claim"

to --Claim 1--

CLAIM 6: Line 1, change "any preceding claim"

to --Claim 1--

CLAIM 7: Line 1, change "any preceding claim"

to --Claim 1--

CLAIM 8: Line 1, change "one of Claims 1 to 5"

to --Claim 1--

CLAIM 9: Line 1, change "any preceding claim"

to --Claim 1--

CLAIM 10: Line 1, change "any preceding claim"

to --Claim 1--

CLAIM 13: Line 1, delete "or 12"

CLAIM 14: Line 1, delete ", 12 or 13"

CLAIM 15: Line 1, change "any of Claims 11 to 14"

to --Claim 11--

CLAIM 16: Line 16, change "any of Claims 10 to 15"

to --Claim 10--

CLAIM 20: Line 1, delete "and 17"

CLAIM 21: Line 1, change "any of Claims 10 to 15"

to --Claim 10--

CLAIM 23: Line 1, delete "and 21"

CLAIM 24: Line 1, change "any of Claims 10 to 15"

to --Claims 10--

CLAIM 26: Line 1, delete "or 25"

CLAIM 27: Line 1, delete "and 24"

CLAIM 28: Line 1, change "one of Claims 10 to 27"

to --Claim 10--

CLAIM 29: Line 1, change "one of Claims 1 to 9"

to --Claim 1--

CLAIM 32: Line 1, change "one of claims 1 to 9"

to --Claim 1--

REMARKS

The specification has been amended to provide a crossreference to the previously filed International Application.

If necessary, the Commissioner is hereby authorized in this, concurrent, and future replies, to charge payment or credit any overpayment to Deposit Account No. 02-2448 for any additional fees required under 37 C.F.R. § 1.16 or under 37 C.F.R. § 1.17; particularly, extension of time fees.

Respectfully submitted,

BIRCH, STEWART, KOLASCH & BIRCH, LLP

P.O. Box 747
Falls Church, VA 22040-0747
(703) 205-8000
Docket No. 0696-0171P

ADM/cw 0696-0171P TPRB

l

A soluble magnesium dihalide complex, its preparation and use

The invention relates to a soluble complex comprising a magnesium dihalide and an electron donor. The invention also relates to a process for the preparation of such a complex, as well as the use of such a complex for the preparation of a polymerization catalyst component containing magnesium, transition metal, halogen and electron donor.

A complex is, according to Römpps Chemie-Lexikon, 7. Edition, Franckh'sche Verlagshandlung, W. Keller & Co., Stuttgart, 1973, page 1831, "a derived name of compounds of higher order, which originates from the combination of molecules, - unlike compounds of first order, in the creation of which atoms participate".

Generally, so called Ziegler-Natta catalyst components have been prepared by reacting a magnesium dihalide-alcohol complex compound or a magnesium alkoxide non-complex compound with a titanium halide and an electron donor which usually is a phthalic acid ester. It is necessary that the magnesium dihalide is amorphous for the catalyst component to be active. Amorphous magnesium dihalide is thus produced in situ.

When reacting the titanium halide with the magnesium dihalide-alcohol complex compound or the magnesium alkoxide non-complex compound, the titanium halide form titanium alkoxy trihalide, which is a harmful waste product. Both reactions have the disadvantage that the titanium halide is wasted for other purposes than the direct provision of catalytically active sites, such as chlorination of the magnesium reactant and washing away of the harmful titanium alkoxy trihalide.

In the production of amorphous magnesium dihalide, such as MgCl₂ strong polar ligand groups (L₁) are needed in order to break up the strong electrostatic crystallinic bonds between the MgCl₂ molecules according to reaction (1):

$$xMgCl2 + xL1 = x(Cl2Mg--L1)$$
 (1)

In practice, polar solvents are needed to carry out reaction (1). In several cases these polar solvents are reactive towards other parts of the catalyst component, and thus have to be replaced by less polar solvents (L_2) . These less polar solvents are, however, often unable to react and co-ordinate with $MgCl_2$ due to the strong intermolecular forces in the $MgCl_2$ structure (2):

5

10

15

20

25

30

15

20

25

30

Typical examples of solvents that are able to form complexes with MgCl₂ are alcohols and water. These compounds have, however, a reactive hydrogen in the hydroxyl group of their molecule which easily reacts with other compounds such as titaniumhalides. Examples of less reactive solvents are the organic esters. They are less reactive towards other components but at the same time they do not have the ability to break up the strongly co-ordinated MgCl₂ molecules. In view of the teaching of the prior art, it seems impossible to achieve amorphous MgCl₂ without harmful side reactions.

EP-A-0 297 076 discloses adducts of magnesium dihalides with alcohols as electron donors for the use as starting materials for the preparation of olefin polymerization catalysts. Alcohols have a reactive hydrogen which react with titanium halides.

The purpose of the invention is therefore to produce amorphous magnesium dihalide in situ without wasting titanium halide or producing harmful waste products. The invention also aims at a stoichiometric route for the preparation of Ziegler-Natta catalyst components and their intermediates. By a stoichiometric route, new catalyst components for the production of tailor-made olefin polymers can be produced.

The purposes of the invention has been achieved by means of a complex comprising a magnesium dihalide and an electron donor, which is characterized in that it is a complex of the magnesium dihalide and the electron donor and has the formula (I):

$$MgX_2 \cdot [R(OR')_n]_m$$
 (I)

wherein MgX₂ is the magnesium dihalide and R(OR')_n is the electron donor, X is a halogen, R is an n-valent C₁-C₂₀ aliphatic group, an n-valent C₇-C₂₇ araliphatic group or an n-valent C₂-C₂₂ acylic group, R' is a C₁-C₂₀ alkyl group or a C₇-C₂₇ aralkyl group, n is a number from 1 to 6 and m is defined as a number $0.5 \le m \le 2.0$. By "n-valent acylic group" is meant a group having n acyl moieties.

Formula (I) is an empirical or semiempirical formula, i.e. m expresses the ratio between the the electron donor R(OR')n and magnesium dihalide MgX_2 . The structural formula may have several molecules of MgX_2 and several same or different molecules of $R(OR')_n$, such as in the complex $(MgX_2')_a \cdot [R(OR')_n]_b$ wherein b:a=m. See e.g. formula (III) below. The claimed complex may be a statistical one, being a mixture of complexes having the average formula (I), or a specific one, essentially all the molecules of which having the same formula (I).

15

In the magnesium dihalide molecular component MgX₂ of the complex, X is preferably selected from Cl, Br and I, and is most preferably Cl. The most preferred complex according to the invention is thus a magnesium dichloride complex.

In the electron donor molecular component $R(OR')_{\Pi}$ of the complex, R is preferably an n-valent C_2 - C_{22} acylic group, more preferably an n-valent aromatic C_7 - C_{22} acylic group, most preferably phthaloyl. R' is preferably a C_6 - C_{16} alkyl, most preferably a C_6 - C_{12} alkyl like undecyl or 2-ethyl-1-hexyl. It means that the phthalate preferably should be an ester of phthalic acid and a longer-chained alcohol. n is preferably about 1 to about 4, preferably about 2, meaning preference for a phthalic acid diester (phthalic acid is dibasic). By "acylic group" is meant the general name for organic acid groups, which form the remainder of carboxylic acids after removing the hydroxyl group.

Thus, the most preferred complex is that of one of the most efficient magnesium compounds and one of the most efficient internal electron donors in the art of Ziegler-Natta catalysts, namely magnesium dichloride and a phthalic acid ester.

In the claimed complex as a whole, m depicts the average ratio between the electron donor molecular component $R(OR')_n$ and the magnesium dihalide molecular component MgX_2 . m is preferably about 0.67 to about 1.0, most preferably about 0.67 or about 1.0. See below, structural formulas (II) and (III).

The complex according to the invention is preferably a magnesium dichloride phthalic acid ester complex having the formula MgCl₂: [C₆H₄(COOR')₂]_m, wherein R' is the same as above and m is from 0.5 to 2.0, most preferably from 0.6 to 1.8.

According to one embodiment of the complex of the invention, the complex is preferably a magnesium dichloride phthalic acid ester complex having the structural formula (II):

$$MgCl_2 \cdot C_6H_4(COOR')_2$$
 (II)

wherein R' is the same as above.

According to another embodiment of the complex of the invention, the complex is a magnesium dichloride phthalic acid ester complex having the structural formula (III):

$$(MgCl2)3 \cdot [C6H4(COOR')2]2$$
 (III)

10

15

20

25

wherein R' is the same as above.

Typically, the claimed complex has an X-ray diffraction pattern (measured by a Siemens D500 instrument equipped with a Cu anode and a graphite monochromator in the reflected beam and using an effect of 40 kV and 35 mA and a CuK α radiation wavelength of 1.541 Å), showing a dominant peak at 4.5° 2 Θ .

The invention also relates to a process for the preparation of a complex comprising a magnesium dihalide and an electron donor.

Characteristic of the claimed process is that a magnesium compound (a) containing an alkoxy moiety, which magnesium compound is selected from the group consisting of a complex of a magnesium dihalide and a magnesium dialkoxide, a complex of a magnesium dihalide and an alcohol, and a non-complex magnesium dialkoxide, is reacted with a halogen compound (b), which is capable of forming the electron donor by replacement of its halogen by said alkoxy moiety.

According to the most important aspect of the invention, the process is a part of a novel stoichiometric preparation process leading to novel catalyst components for olefin polymerization.

The halogen compound (b) is capable of forming the electron donor by replacement of its halogen by said alkoxy moiety. This does not only mean that the halogen compound is a reagent leading to the electron donor, but also that it is a structural precursor thereof. Compare e.g. an alkyl halide with the corresponding dialkyl ether or an organic acid halide with the corresponding organic acid ester.

The halogen compound (b) preferably has the formula (IV):

RXn (IV)

wherein R is an n-valent C_1 - C_{20} aliphatic group, an n-valent C_7 - C_{27} araliphatic group or an n-valent C_2 - C_{22} acylic group, X is a halogen and n is 1 to 6. In the formula, R is preferably an n-valent C_2 - C_{22} acylic group, more preferably an n-valent aromatic C_7 - C_{22} acylic group, most preferably phthaloyl. X is preferably selected from Cl, Br and I, and is preferably Cl. n is preferably 1 to 4, most preferably about 2.

According to a preferred embodiment of the process of the invention, said halogen compound (b) is an organic acid halide, preferably phthalic acid dichloride Ph(COCl)2, wherein Ph is o-phenylene. When one considers that the most preferred

10

15

20

25

30

internal electron donor molecule of the claimed complex is a phthalic acid ester, the complexed ester molecule is simply formed by replacing the chlorines of the phthalic acid dichloride with alkoxy groups.

According to a first alternative embodiment of the process of the invention, said magnesium compound (a) containing an alkoxy moiety is a magnesium dichloride-magnesium dialkoxide complex of the formula (V):

$$\label{eq:mgCl2} \text{MgCl}_2 \cdot [\text{Mg(OR')}_2]_p \tag{V}$$

wherein R' is a C_1 - C_{20} alkyl group or a C_7 - C_{27} aralkyl group, preferably a C_6 - C_{16} alkyl group, and p is 1 to 6, preferably about 2. Formula (V) is empirical or semiempirical, meaning that the complex's molecular clusters can consist of several $MgCl_2$ molecules and several $Mg(OR')_2$ molecules, corresponding to the formula $(MgCl_2)_c[Mg(OR')_2]_d$ wherein d:c = p. In the process of the invention, said magnesium dichloride-magnesium dialkoxide complex is reacted with the above halogen compound so that the halogen compound (b) has its halogen(s) replaced by the alkoxide(s) of the complex and forms an electron donor, whereby the complex of magnesium dichloride and the electron donor is formed.

Preferably, said complex of a magnesium dialkoxide and a magnesium dialkoxide is a magnesium dichloride-dimagnesium dialkoxide complex of the structural formula (VI):

$$MgCl_2 \cdot [Mg(OR')_2]_2$$
 (VI)

wherein R' is a C_1 - C_{20} alkyl group or a C_7 - C_{27} aralkyl group, preferably a C_6 - C_{16} alkyl group.

Said magnesium dichloride-magnesium dialkoxide complex is preferably prepared by reacting magnesium dichloride and an alcohol into an intermediate which is a magnesium dichloride-alcohol complex $MgCl_2 \cdot (R'OH)_{2p}$ wherein R' is the same as above, and reacting the magnesium dichloride alcohol complex with p mol of a magnesium dialkyl $MgR"_2$, wherein R" is a hydrocarbyl group having 1 to 20 carbon atoms. If R" is a hydrocarbyl group having 1 to 5 carbon atoms, a volatile alkane R"H byproduct is thereby formed and easily removed by evaporation. In the synthesis, the molar ratio $MgCl_2:R'OH$ is preferably between 1:1 and 1:8, most preferably between 1:2 and 1:5. The molar ratio $MgCl_2:(R'OH)_{2p}:MgR"_2$ is preferably between 1:1 and 1:4, most preferably about 1:2.

25

Said magnesium compound (a) which is said magnesium dichloride-dimagnesium dialkoxide complex $MgCl_2 \cdot [Mg(OR')_2]_2$, wherein R' is the same as above, is preferably reacted with said halogen compound (b) which is said phthalic acid dichloride $Ph(COCl)_2$, wherein Ph is o-phenylene. Typically, the product is $(MgCl_2)_3 \cdot [Ph(COOR')]_2$. See formula (III) above.

In the first alternative embodiment of the claimed process, said magnesium compound (a) and said halogen compound (b) are reacted in essentially stoichiometrical amounts and, independently, at a temperature of between 80 °C and 160 °C. The reaction time is preferably about 2 h to about 8 h.

- Most preferably, the magnesium dihalide and the alcohol, which is a heavier alcohol, are first reacted at a temperature between 120 °C and 160 °C, after which the product is reacted with the magnesium alkoxide at a temperature between 80 °C and 120 °C, followed by reaction with the halogen compound at said lower temperature interval.
- 15 A typical example of said first alternative embodiment of the claimed process is described in Figure 1.

According to a second alternative embodiment of the claimed process, said magnesium compound (a) containing an alkoxy moiety is a non-complex magnesium dialkoxide of the formula (VII):

 $Mg(OR')_2 (VII)$

wherein R' is a C_1 - C_{20} aralkyl group or a C_7 - C_{27} aralkyl group, preferably a C_6 - C_{16} alkyl group.

Said non-complex magnesium dialkoxide is preferably prepared by reacting a magnesium dialkyl, preferably a magnesium dialkyl of the formula MgR"₂, wherein R" is a hydrocarbyl group having 1 to 20 carbon atoms, and an alcohol, preferably an alcohol of the formula R'OH, wherein R' is the same as above. The preferred ratio between the magnesium dialkyl and the alcohol is about 1:2.

In the second alternative embodiment of the claimed process, said magnesium compound (a) which is said non-complex magnesium dialkoxide, has the formula Mg(OR')2, wherein R' is a C₁-C₂₀ aralkyl or a C₇-C₂₇ aralkyl, preferably a C₆-C₁₆ alkyl, is preferably reacted with said halogen compound (b) which is said phthalic acid dichloride Ph(COCl)₂, wherein Ph is o-phenylene. Usually, said

20

25

30

magnesium compound (a) and said halogen compound (b) are reacted in essentially stoichiometric amounts. The product formed is preferably MgCl₂·Ph(COOR')₂. See Formula II above.

A typical example of said second alternative embodiment of the claimed process is described in Figure 2.

According to a third alternative embodiment of the process of the invention, said magnesium compound (a) containing an alkoxy moiety is a complex of a magnesium dihalide and an alcohol of the Formula (VIII):

$$MgCl_2 \cdot (R'OH)_{q}$$
 (VIII)

wherein R' is a C₁-C₂₀ alkyl group or a C₇-C₂₇ aralkyl group, preferably a C₆-C₁₆ alkyl group, and q is from 1 and 6. The alkoxy moiety is the R'O group of the alcohol R'OH. This complex is often used as starting material for Ziegler-Natta catalyst components. However, it is not known to have been used as starting material for a magnesium dihalide electron donor complex by reacting it with a halogenous electron donor precursor in the above described way.

The complex of a magnesium dihalide and an alcohol is usually prepared by reacting magnesium dichloride MgCl₂ and and alcohol R'OH, wherein R' is the same as above.

In the process according to the second alternative embodiment, said magnesium compound (a) which is said complex of a magnesium dihalide and an alcohol having the formula $MgCl_2 \cdot (R'OH)_q$, wherein R' is a C_1 - C_{20} alkyl or a C_7 - C_{27} aralkyl, preferably a C_6 - C_{16} alkyl, and q is from 1 and 6, is preferably reacted with said halogen compound (b) which is said phthalic acid dichloride $Ph(COCl)_2$, wherein Ph is o-phenylene. An example of this third alternative embodiment is described in Figure 3.

Above, the product and its preparation according to the invention have been described. As the claimed complex finds its natural application in the field of olefin polymerization catalyst synthesis, the invention also relates to the use of said complex in that field. Thus, claimed is the use of said complex for the preparation of a polymerization catalyst component containing magnesium, as well as at least one transition metal, halogen and electron donor. More specifically, the use is characterized in that said complex is reacted with a titanium halide (c) to give said catalyst component.

Said titanium halide (c) preferably has the Formula (IX):

$$(OR''')_{\mathbf{p}} TiX_{\mathbf{4-p}} \tag{IX}$$

wherein R'" is a C_1 - C_{10} alkyl group or a C_7 - C_{16} aralkyl group, X is a halogen and p is 0 to 3. Most preferably, said titanium halide (c) is a titanium tetrahalide TiX_4 , wherein X is the same as above, most preferably titanium tetrachloride $TiCl_4$.

Experimental

5

15

20

25

30

Preparation of the complexes

All chemicals were handled in strict inert conditions and all the reactions took place also in strict inert conditions in a nitrogen atmosphere.

10 Example 1 (first alternative embodiment)

First 1.07 g (11.2 mmol) of MgCl₂ was introduced into a 50 ml glass reactor. 9.60 ml (8.0 g, 46.2 mmol) of 1-undecanol was added on to the MgCl₂. The slurry was mixed using a magnetic stirring bar and the solution was heated to 130 °C and the reactants were allowed to react with each other at this temperature for 3 h. The slurry was cooled down to 100 °C and 9.6 ml (8.3 g, 90 mmol) of toluene was added to the reaction solution to increase its dissolving capability. 25.40 ml (18.52 g, 22.3 mmol R"₂Mg) of a 20% heptane solution of butyl-octyl magnesium was now introduced. Finally, 3.24 ml (4.565 g, 22.5 mmol) of phthaloyl chloride was added. The achieved product was dried under a stream of nitrogen for several hours at temperatures between 90 °C and 120 °C.

Example 2 (second alternative embodiment)

123.2 mmol of butyl-octyl magnesium was introduced into a 250 ml glass reactor. A 20% heptane solution of the butyl-octyl magnesium containing 2.92 w-% of Mg was used giving a feed volume of 139.4 ml (102.5 g) in the reactor. 244.6 mmol (38.2 ml, 31.85 g) of 2-ethyl-l-hexanol was then added slowly at room temperature. The addition of alcohol took 23 min. Mixing speed was about 240 rpm. The temperature was increased to 63 °C and the reactants were allowed to react with each other at that temperature for 15 min. After this 122.74 mmol (17.69 ml, 24.92 g) of phthaloyl chloride was added slowly at room temperature. The temperature was increased during 10 min to 50 °C and the reactants were again allowed to react with each other at that temperature for 5 min. After this the reaction solution was allowed to cool down to room temperature.

10

15

20

28.8 g of the achieved solution was taken into a 100 ml glass reactor for solvent evaporation. The sample was dried under vacuum in a stream of nitrogen gas at 50 °C for 3 h. 12 ml of condensed solvent (heptane) was trapped in the vacuum trap.

The product was washed with 60 ml of pentane at 45 °C for 45 min, after which the product was allowed to settle for 45 min and the solid product was separated from the solution. The product was washed a second time with 44 ml of pentane and finally dried under vacuum and in a stream of nitrogen at 50 °C for one hour.

Example 3 (third alternative embodiment)

22.60 mmol (2.15 g) of MgCl₂ was introduced into a 100 ml glass reactor. To this, 45.19 mmol (7.10 ml, 5.92 g) of 2-ethyl-1-hexanol was added. Finally, 22.60 mmol (3.26 ml, 4.59 g) of phthaloyl dichloride was added to the mixture. The mixture was kept under agitation at 60 °C for 30 min. A solid complex was collected from the vessel by evaporation and washed three times with a 100 ml portion of heptane at 90 °C for 15 min, then with a 100 ml portion of pentane at room temperature and finally dried. This product was reacted with TiCl₄ into a catalytically active complex.

Comparative example

A fourth sample was prepared by introducing 20 mmol MgCl₂ (1.90 g) in a 150 ml glass reactor equipped with a magnetic stirrer. 20 mmol of di-2-ethyl-hexyl-phthalate (8.0 ml, 7.81 g) (DOP) was then added on to the MgCl₂. The reactants were allowed to react with each other overnight. The product was washed with pentane and dried in the same way as described above.

Characterization of the products by X-ray diffractometry and Infrared spectroscopy

The products were characterized by infrared spectroscopy (IR) and by taking X-ray diffraction patterns of it. The WAXS patterns were collected in reflection mode between 2 and 70°2 θ with a Siemens D500 instrument. The diffractometer was equipped with a Cu anode and a graphite monochromator in the reflected beam. The effect used was 40 kV and 35 mA. The CuKα radiation wavelength was 1.541 Å.
 The sample was loaded in a glove box into a Mylar film covered sample holder.

20

25

Di-undecylphthalate (DUP) was used as electron donor reference in the IR studies. The products of examples 1 and 2 were investigated together with the standard DUP. The products of example 1 was analyzed twice: right away and a second time after overnight storage.

The IR spectra were taken by a Nicolet 510 FTIR equipment with 2 cm⁻¹ resolution. The number of scans were 128. All the samples were investigated as capillary films between two KBr pellets. Pure DUP was not handled in inert conditions, but the MgCl₂ samples were handled in a glove box in an inert nitrogen environment in order to protect the samples from air and moisture. To get thin enough capillary films the samples were somewhat heated when placed in between the KBr pellets.

X-ray characterization

The X-ray diffraction pattern of the MgCl₂-DOP complex is shown in Figure 4. The pattern show no sign of MgCl₂. At 50 ° 2 Θ , where the most significant peak of pure MgCl₂ is to be found, there is no sign of a peak. This is also the case at 30° and 35° 2 Θ . On the other hand, there is a new dominant peak at 4.5° 2 Θ showing that the reflecting layers have been separated far from each other (21 Å).

The X-ray diffraction pattern of the product coming from the direct reaction between $MgCl_2$ and DOP of the comparative example is shown in Figure 5. The pattern showed that no reaction between these components had occurred, the X-ray pattern showed pure $MgCl_2$ with its significant reflecting signals at 15°, 30°, 35° and 50° 2Θ .

The results show that in the first three synthesis routes the donor compound had been complexed with $MgCl_2$ at a molecular level, thereby breaking up the strong molecular structure of crystalline $MgCl_2$ and indicating in situ preparation thereof. The results show also that in the fourth, direct synthesis route, the donor compound had not been able to form a complex with $MgCl_2$ but the product contained only original $MgCl_2$.

Results by Infrared Spectroscopy

The samples were prepared and studied by IR spectroscopy according to the description in the experimental section above. As MgCl₂ does not have any absorbance spectrum in the IR area from 4000 to 400 cm⁻¹, the IR study concentrated on the changes in the structure of the DOP and DUP complexation to the MgCl₂.

The pure DOP and DUP absorbs IR light due to the stretching vibration of the ester carbonyl double bond of the C=O group at the wavelength of 1729 cm⁻¹. The corresponding stretching vibration of the C-O- bond is to be found at 1280 cm⁻¹ and at 1100 cm⁻¹ (see Figure 6).

- When the MgCl₂ was co-ordinated to DOP in the molar ratio of 1:1, clear changes in the IR spectrum of DOP could be seen (figure 6, Example 2). The absorption peaks of pure DOP could still clearly be seen which indicates that a part of the carbonyl groups are still unco-ordinated. On the right side of the original C=O absorption peak there is a new shoulder indicating that a part of the C=O groups have co-ordinated to Mg causing the double bond of the C=O group to loosen up starting to resemble more a single C-O bond. The proportionally small shift in the position of the "shoulder peak" indicate a weak interaction, i.e. a weak co-ordination of MgCl₂ to the C=O oxygen. A weak interaction is also indicated by the sign of several secondary "shoulder" peaks.
- When the MgCl₂ amount was increased by 50% in the complex (Example 1), the proportion of the "shoulder" peak at about 1690 cm⁻¹ increased. There is however no sign of a strong co-ordination (figure 6). The changes in the IR spectra show up even better in the sample that had been stored overnight. Here the main peak of the carbonyl oxygen has shifted from the position of 1729 cm⁻¹ for the pure DUP to 1719 cm⁻¹. The results show also that the co-ordinated carboxyl group is influencing the "free" carboxyl group as the position of its peak is shifted 10 cm⁻¹.

The same results can be seen in connection with the absorption peak of the C-O-bond. The absorption peak of the C-O-bond in the pure DUP is found at 1287 cm⁻¹. Looking at the spectra for the MgCl₂·DOP (Example 2) and the (MgCl₂)_{1.5}·DUP (Example 1) samples there is a corresponding "shoulder" peak forming to the left of the original peak indicating a weak double bond character of the C-O bond. This shift is so strong that the original peak at 1287⁻¹ is not any longer detectable in the product that had been stored overnight (figure 6). These results indicate that the Mg in the MgCl₂ is complexed between the C=O oxygen and the C-O oxygen atoms in the MgCl₂·DOP and in the MgCl₂·DUP complex.

Conclusively it can be said that in the IR spectrum of the pure DUP the peaks show unco-ordinated carbonyl groups, in the $MgCl_2 \cdot DOP$ and in the $MgCl_2 \cdot DUP$ sample (Example 2) there is a $MgCl_2$ co-ordination to one of the carbonyl groups, the other being free, and in the $(MgCl_2)_{1.5} \cdot DUP$ sample (Example 1) there is a partial co-ordination of $MgCl_2$ to both of the carbonyl groups.

10

15

20

35

Examples 4 to 7 (use of the complex according to the first alternative embodiment)

Preparation of the catalyst component complex

1.69 g (17.70 mmol) of anhydrous MgCl₂ was introduced in inert conditions into a 100 ml septum bottle. 11.12 ml (9.27 g, 70.80 mmol) of 2-ethyl-hexanol (EHA) was introduced on to the MgCl₂ and after this the temperature was increased to 125-128 °C to allow the reaction components to react with each other. After this, 8.81 ml (7.67 g, 83.19 mmol) of toluene was added after the reaction solution had cooled down to 110 °C. After the addition of the toluene the reaction solution was cooled down to 21 °C. Then 40 ml (29.16 g, 35.4 mmol) of a 20 w-% heptane solution of butyl-octyl-magnesium (BOMAG) was added. After this 5.10 ml (7.19 g, 35.4 mmol) of phthaloyl dichloride (PDC) was added to produce a MgCl₂ donor complex solution.

Use of the catalyst component complex

The MgCl₂ donor complex according to the first alternative embodiment was now, drop by drop, added into 38.91 ml (67.16 g, 354 mmol) of TiCl₄ and allowed to react with this reagent at a temperature of 95 °C. The reactants were allowed to react with each other for 30 min.

After the TiCl4 treatment, the complex was allowed to settle and the liquid was siphoned off. After this, 100 ml (86.6 g, 0.94 mol) of toluene was added on to the complex and the complex was washed in this solution at 90 °C for 20 min. Depending on which of the synthesis was under work, this washing step was done once (Example 1), twice (Example 2), three times (Example 3) or four times (Example 4). Finally, the catalyst complex was washed twice with 65 ml (44.44 g, 0.44 mol) portions of heptane for 20 min at 80 °C and thereafter, the complex was washed at room temperature with a 55 ml (34.44 g, 0.48 mol) portion of pentane for 20 min to improve the drying conditions. The catalysts were dried under a stream of nitrogen for one hour.

Chemical characterization of the complexes

The catalyst complexes were characterized with respect to their chemical composition by measuring their Ti and Cl content. The Ti analysis was started by

20

25

30

35

dissolving the samples in a mixture of nitric and hydrofluoric acid. The metal was measured flame atomic absorption with a nitrous acetylene flame. Chloride was determined after dissolution in dilute sulphuric acid by potentiometric titration with a standard silver nitrate solution.

5 Determination of donors and phthalic anhydride

The determination of the phthalic esters and the phthalic anhydride were done by first dissolving the sample in acetone. The dissolving was improved by keeping the acetone slurry in an ultra-sound bath for 5 min. After this the samples were filtered and run by solution chromatography. As eluent a solution consisting of water and acetonitrile in the proportion of 4/96 was used. Eluent flow rate was 1.5 ml/min. A photo diode array was used as detector. Each component was identified by comparing the respective retention time and UV spectra with standard components.

15 GC studies to measure alcohol content

To check the conversion rate of the ethanol (EtOH), 2-ethyl-hexanol (EHA), or other alcohol added in the synthesis, the alcohol content of the catalysts were measured by gas chromatography (GC). This was done by first dissolving a 100 mg sample of the catalyst in 1 ml of n-pentanol. Depending on the alcohol to be measured, an internal alcohol standard was chosen. If ethanol was to be measured the n-pentanol solution contained n-propenol as internal standard. To improve the solubility of the catalyst in the solution, the sample was kept in an ultra-sound bath. To remove the inorganics from the organic solution it was extracted with 1 ml of water and to ensure full dissolution, another ml of the n-pentanol solution was added. To ensure repeatable equilibrium conditions between the organic layer and the water layer the samples were allowed to stand overnight. The sample for the GC was taken from the alcohol layer. A Hewlett Packard 5890 GC with a 60 m DB-1 column was used for the GC analyses. The column had a diameter of 0.25 mm with a film thickness of 1 μm. An FID detector was used.

Bulk polymerization

Propylene was polymerized in stirred tank reactor having a volume of 5 l. About 0.9 ml triethyl aluminium (TEA) as a cocatalyst, ca 0,12 ml of a 100-% solution of cyclohexyl methyl dimethoxy silane as an external donor and 30 ml of n-pentane were mixed and allowed to react for 5 minutes. Half of the mixture was added to the

polymerization reactor and the other half was mixed with ca 20 mg of the catalyst complex. After additional 5 minutes the catalyst/TEA/donor/n-heptane mixture was introduced into the reactor. The Al/Ti mole ratio was 250 and the Al/external donor mol ratio was 10 mol/mol. 70 mmol hydrogen and 1400 g of propylene were introduced into the reactor and the temperature was raised within 15-30 minutes to 70 °C. The polymerization time was 60 minutes, after which the polymer formed was taken out from the reactor. The polymers were characterized with respect to their Melt Flow Rate (MFR2), bulk density (BD) and fraction of total solubles in xylene (TS).

10

15

5

Results

Preparation of the complexes

The catalyst complexes achieved in this investigation are listed in Table 1.

Table 1

The catalyst complexes prepared.

Example	Number of toluene washes	Colour of catalyst	Morphology of catalyst
4	1	Dark wine-red	Freely flowing
5	2	Dark wine-red	Freely flowing
6	3	Dark wine-red	Freely flowing
7	4	Dark wine-red	Freely flowing

20

25

The chemical composition of the catalysts

The chemical composition of the catalysts were measured according to the description in the experimental section. In Table 2 the chemical composition of the catalysts are listed in w-% units, in Table 3 the composition is listed in mol-% units and in Table 4 the molar proportions between Mg, Ti and DOP are compared.

30

The chemical compositions of the catalysts were as expected on the basis of the reaction equation. With three washes a composition of (MgCl₂)₆TiCl₄DOP was achieved. During the washes, there was a slightly higher wash out of TiCl4 compared to DOP in the last catalyst. The amount of free alcohol (EHA) was also

very low playing no significant part in the chemical composition (now 0.004-0.006 mol-%), i.e. being about 5% of the mol amount of TiCl₄ or DOP. The amount of phthalic anhydride was about 50% of the DOP amount. To sum up the results from the chemical measurements it can be said that the chemical composition of the catalyst complex when using the MgCl₂ enriched Mg(OR')₂ as a reagent in the catalyst synthesis is (MgCl₂)₃TiCl₄DOP(PA)_{0.5}.

Table 2

The chemical composition of the catalysts in w-% units

Example	Mg w-%	Ti w-%	DOP w-%	EHA w-%	PA w-%
4	7.8	4.7	33.6	0.72	6.7
5	8.1	4.7	32.0	0.54	7.4
6	10.2	3.2	28.5	0.58	6.5
7	12.9	1.6	21.6	0.51	6.1

Table 3

The chemical composition of the catalysts in mol-% units

Example	Mg	Ti	DOP	EHA	PA
	mol-%	mol-%	mol-%	mol-%	mol-%
4	0.321	0.098	0.086	0.0055	0.045
5	0.333	0.098	0.082	0.0041	0.050
6	0.420	0.067	0.073	0.0045	0.044
7	0.531	0.033	0.056	0.0039	0.041

Table 4

The molar ratio between Mg, Ti and DOP

Example	Mg	Ti	DOP
4	3.3	1	0.9
5	3.4	1	0.8
6	6.3	1	1.1
7	15.9	1	1.7

Calculated and found chlorine contents

The chlorine content in the catalysts were calculated on the basis of the Mg and Ti content. The calculations were based on the assumption the Mg was present in the catalyst as MgCl₂ and Ti as TiCl₄. These calculated results were then compared to the measured results. The results are listed in Table 5. The results showed to be in good agreement, which indicates that both Mg and Ti are present in the catalyst complexes in the fully chlorinated form.

10 Table 5

5

The calculated and the found chlorine content in the catalysts

Example	Calculated Cl w-%	Found Cl w-%
4	36.7	36.9
5	37.6	38.0
6	39.3	39.7
7	42.4	43.8

Wash out of TiCl4.DOP

15

All the chemical measurements support the same conclusion: due to the toluene, TiCl₄ and DOP are washed out from the catalyst in a molar proportion of 1:1. This shows up as a constant decrease of the Ti mol-% and the DOP mol-%, and as a constant increase of the Mg mol-% and the Cl mol-%.

20

25

Activity of the catalysts

All the catalyst complexes were test polymerized according to the descriptions in the experimental section. The results are listed in Table 5. The results showed that all the catalyst complexes had about the same activity, being between 1.0 and 1.5 kg PP/g cat.

Table 6

The test polymerization results

Example	Activity kg PP/g cat	Activity kg PP/g Ti
4	1.1	23
5	1.2	26
6	1.5	45
7	1.3	81

5 MFR of the polymers

Table 7

In Table 7 the MFR values achieved from the test polymerization results are listed. The results indicated a systematic increase in MFR with increasing number of toluene washes as MFR increases from 2.0 in the first polymer to 13.7 in the third.

The MFR values of the polymers

Example	MFR
4	2.0
5	4.9
6	13.7
7	12.4

15 Examples 8 to 13 (use of the complex according to the second alternative embodiment)

Preparation of the catalyst component complexes

- 20 All chemicals were handled in strict inert conditions and all the reactions took place also in strict inert conditions in nitrogen atmosphere.
- 8.85 mmol of butyl-octyl-magnesium was introduced into a 150 ml glass reactor. A 20% heptane solution (BOMAG-A) was used giving a feed volume of 10 ml (7.29 g). 17.7 mmol (2.78 ml, 2.32 g) of 2-ethyl-1-hexanol (EHA) was then added at room temperature. The temperature was increased to 60 °C and the reactants were

10

15

20

25

30

35

allowed to react with each other at that temperature for 30 min. After this 8.85 mmol (1.28 ml, 1.80 g) of phthaloyl chloride (PDC) was added and the reactants were again allowed to react with each other for 30 min at 60 °C to give the claimed complex.

The resulting solution of the claimed complex was added dropvise into 88.5 mmol (9.73 ml, 16.79 g) of TiCl₄ that had been preheated to 95 °C. The reactants were also in this case allowed to react with each other for 30 min at 95 °C. After this 60 ml of toluene was added. After the precipitate had settled the mother liquid was siphoned off. Five different examples were carried out according to this description. After this the catalyst complex was washed with 30 ml portions of toluene. In Example 8, the complex was washed once with toluene, in Example 9 twice, in Example 10 three times, in Example 11 four times and in Example 12 six times with 30 ml portions of toluene. The toluene washes were carried out at 90 °C. Finally, the complex was washed three times with 30 ml portion of pentane. The complexes were finally dried under a stream of nitrogen. The yield of the catalyst was about 2 g which corresponded to about 75% of the theoretical.

Characterization of the catalyst components

The catalyst component complexes were analyzed with respect to their Mg, Cl and Ti content. In addition to this, the amount of donor compound, the di-octyl-phthalate (DOP) formed in the synthesis, was measured from the catalysts. To indicate to what degree the formed donor compound (DOP) was decomposing in the synthesis, the amount of phthalic anhydride (PA) was also measured from the catalysts.

IR and X-ray of the unwashed Mg:Ti:DOP complex

A stoichiometric complex of MgCl₂·TiCl₄·DOP was prepared by reacting 6.37 mmol (7.19 ml, 5.24 g) of BOMAG with 12.73 mmol (2.00 ml, 1.67 g) of EHA in a 50 ml glass reactor. After this 6.365 mmol (0.92 ml, 1.29 g) of phthaloyl chloride was introduced and last 6.37 mmol (0.70 ml, 1.21 g) of TiCl₄ was added. The solid product was washed with pentane and finally, the sample was dried in a stream of nitrogen. The sample was characterized by IR spectroscopy and by means of its X-ray diffraction pattern.

The IR studies

IR spectres were taken by means of a Nicolet 510 FTIR equipment with 2 cm⁻¹ resolution. The number of scans were 128. All the samples were investigated as capillary films between two KBr tablets. The pure EHA was not handled in inert conditions, while the MgCl₂ samples were handled in a glovebox in an inert nitrogen environment in order to protect the samples from air and moisture.

X-ray diffraction patterns

15

10

The WAXS patterns were collected in a reflection mode between 2° and 70° 2Θ with a Siemens D500 instrument. The diffractometer was equipped with a Cu anode and a graphite monochromator in the reflected beam. The CuK α radiation wavelength was 1.541 Å. The effect used was 40 kV and 35 mA. The sample was loaded in a glovebox into a Mylar film covered sample holder.

Bulk polymerization

The bulk test polymerization was carried out according to the above description relating to the first alternative embodiment.

20

25

Results

Preparation of the complexes

The reaction between the Mg-alkyl and the alcohol resulted in a clear solution with a little bit higher viscosity. The reaction was exothermic as the solution became warm when mixing the reactants, the temperature increase was from room temperature up to 50 °C. When the phthaloyl chloride was added a slight yellow colour appeared. Also this reaction was slightly exothermic. The reaction solution become again freely flowing with a low viscosity.

The TiCl₄ was introduced into a 150 ml glass reactor and heated to 95 °C. The Mg solution was then added to the hot TiCl₄ solution dropwise. A beige precipitate started to form right at the beginning of the addition. During addition the solution turned turbid. A partly freely floating precipitate was formed together with more tarlike precipitate that started to foal the reactor walls. To improve the settling conditions toluene was added to the reaction solution. A satisfactory settling of the product was then achieved so that the reaction solution could be siphoned off.

Depending on the number of toluene washes the resulting product become more freely flowing. If only one toluene wash was used the product was still as agglomerates, but already two toluene washes resulted in a freely flowing powder-like product.

In the case of the catalyst components of examples 5, 6 and 7, a joined MgCl₂-DOP complex and a joined addition to the TiCl₄ solution was carried out. After the first toluene wash, 1/3 of the solution slurry was separated. The separated part was then washed with the aliphatic hydrocarbon and dried to give the product of example 5. The remaining part of the slurry was washed a second time with toluene and half of this solution slurry was then taken out from the reactor and undertaken the same hydrocarbon treatment as in Example 5, resulting in the product of Example 6. The remaining part of the catalyst slurry in the reactor was washed twice with toluene and then washed with an aliphatic hydrocarbon in the same way as the first two examples. This sample was the product of example 7. The catalyst morphologies are listed in Table 8.

Table 8

The morphology of the catalysts

Example	Number of toluene washes	Morphology of catalyst
8	1	Black agglomerates
9	2	Dark powder
10	4	Dark powder

The Chemical composition of the catalysts

The Mg, Ti, Cl, DOP, EHA and the phthalic anhydride (PA) content of the catalysts were measured. The results are listed in w-% units in Table 9. In Table 10 the chemical composition is given in mol-% units and in Table 11 the Mg and DOP amounts are compared to the Ti amount on a molar basis. Table 12 shows the Cl content of the catalysts.

2.0

25

5

10

15

Table 11

Table 12

Table 9

The chemical composition of the catalysts in w-% units

Example	Mg w-%	Ti w-%	DOP w-%	EHA w-%	PA w-%
8	5.7	6.8	47.7	0.26	3.4
9	11.3	3.1	32.2	0.18	2.5
10	13.4	1.4	21.3	0.25	1.7

Table 10

The chemical composition of the catalysts in mol-% units

Example	Mg mol-%	Ti mol-%	DOP mol-%	EHA mol-%	PA mol-%
8	0.235	0.142	0.122	0.002	0.023
9	0.465	0.065	0.083	0.001	0.017
10	0.551	0.029	0.055	0.002	0.012

The molar proportions between Mg and Ti and between DOP and Ti

Example	Mg	Ti	DOP
8	1.7	1	0.86
9	7.2	1	1.28
10	18.9	1	1.87

The calculated Cl content in the catalysts compared to the measured amounts

Example	Calculated Cl w-%	Found Cl w-%
8	36.8	36.4
9	42.2	45.0
10	43.3	44.5

Activity of the catalysts

All the catalysts were test polymerized according to the above instructions. The polymerization results are listed in Table 13 in both kg PP/g cat and kg PP/g Ti units. Activities of almost 8 kg PP/g cat were achieved. Catalysis of the Examples 8, 9 and 10 gave good polymerization results, with the highest activity achieved for the catalyst that had been twice washed with toluene. The activities expressed in kg PP/g Ti units showed an linear increase related to the number of toluene washes for the catalysts of Examples 8, 9 and 10. Activities of over 500 kg PP/g Ti were reached.

Characterization of the polymers

All the polymers were characterized with respect to their melt flow rate (MFR) and bulk density (BD). All the polymers showed to have a MFR₂ between 11-12 g/10 min, indicating a quite good hydrogen response. Bulk densities were between 0.350-0.390 g/ml. The total solubles were between 2 and 3%, being better for the polymers achieved with the catalyst giving higher activity. The results listed in Table 14.

20

10

15

Table 13

The polymerization results

Example	Activity kg PP/g cat	Activity kg PP/g Ti
8	2.56	38
9	7.88	254
10	7.33	524

25

Table 14

The polymer properties

Example	MFR 2.16 kg, 10 min	TS %	BD g/ml
8	11.0	3.1	0.360
9	12.37	2.1	0.350
10	11.0	2.1	0.390

IR studies of the catalyst obtained

IR spectra in the corresponding regions of 1500-1950 cm⁻¹ and of 1000-1450 cm⁻¹ were taken from the resulting catalyst of example 9 and compared to an IR spectrum of a typical active catalyst complex coming from a synthesis starting from a MgCl₂-(EtOH)₃ support material. The spectra are essentially different, and also different from the IR spectra of the isolated complexes of TiCl₄/DOP and MgCl₂/DOP.

X-ray studies of the catalyst obtained

As described above, X-ray diffraction patterns were taken from the resulting catalysts and compared to a X-ray pattern from an inactive catalyst complex and a typical active catalyst complex prepared from a MgCl₂·3EtOH support material.

In the X-ray patterns of Mg(OR)₂, MgCl₂·TiCl₄·DOP produced from MgCl₂·3EtOH, and of (MgCl₂)_{1.7}·TiCl₄·DOP produced by adding one mol MgCl₂·DOP to 10 moles of TiCl₄, there was a strong peak located between 5° and 9° 2Θ. In addition, there is a halo formation between 17° and 23° 2Θ. The strong peak in the left corner of the pattern indicates that large organic groups are separating metal layers at a distance of between 9 and 17 Å, the distance depending on the size of the organic compound (DOP or di-undecyl phthalate DUP). It can thus be stated that the X-ray diffraction patterns for the final catalyst complexes originating from the claimed complexes all show unique features originating from the starting compounds of Mg(OR)₂ and MgCl₂·DOP. These patterns show almost no sign of amorphous or crystalline MgCl₂.

Examples 11 (use of the complex according to the second alternative embodiment) and 12 and 13 (use of the complex according to the third alternative embodiment)

The following reagents are used; MgCl₂ or MgR₂, 2-ethyl-hexanol (EHA), phthaloyl dichloride (PDC) and TiCl₄ and they are added in the molecular proportion of 1:2:1:1. In the first synthesis (Example 11), the Mg-alkyl is reacted with the alcohol, then the phthaloyl chloride (PDC) is added and finally the TiCl₄ is added. In the next two syntheses (Examples 12 and 13), the Mg-alkyl is replaced by MgCl₂. Either the TiCl₄ or the phthaloyl chloride is added in the next step, followed by the last reagent. The synthesis set-ups are is listed in Table 16.

30

35

5

10

15

20

25

Table 16

Addition order of the reaction components in the catalyst synthesis

Reaction component/Example	11	12	13
MgR ₂	11		
MgCl ₂	-	1	1
R'OH	2	2	2
PDC	3	3	4
TiCl ₄	4	4	3

Preparation of the complexes

The same volumes of reagents have been used in all the experiments regardless in which order they have been added. Thus 22.22 mmol (25.10 ml, 18.3 g) of a 20% heptane solution of butyl-octyl-Mg (BOMAG) was added in experiments (8) and (9) and 22.60 mmol (2.15 g) of MgCl₂ was added in experiment (10). To this, 45.19 mmol (7.10 ml, 5.92 g) of 2-ethyl-1-hexanol EHA was added. The TiCl₄ mol amount added was equal to the mol amount of MgCl₂ being 22.60 mmol (2.48 ml, 4.29 g) and also equal to the mol amount of PDC added, which was 22.60 mmol (3.26 ml, 4.59 g). The addition orders of the reaction components in each catalyst synthesis are listed in Table 16. All the complexes were washed three times with a 100 ml portion of heptane at 90 °C for 15 min and last with a 100 ml portion of pentane at room temperature. Finally the catalysts were dried under a stream of nitrogen.

Characterization of the catalysts

All the catalysts were characterized with respect to their chemical composition by measuring their Mg, Ti, Cl and di-octyl-phthalate (DOP) content. The Ti and Mg containing catalyst samples were dissolved in a mixture of nitric and hydrofluoric acid and the metals were measured by flame atomic absorption with a nitrous oxide/acetylene flame. Chloride was determined after dissolution in dilute sulphuric acid by potentiometric titration with a standard silver nitrate solution.

The determination of the phthalic esters and the phthalic anhydride were done by first dissolving the sample in acetone. The dissolution was improved by keeping the

20

25

15

5

10

acetone slurry in an ultra-sound bath for 5 min. After this the samples were filtered and run by solution chromatography. As eluent a solution consisting of water and acetonitrile in a proportion of 4/96 was used. The eluent flow rate was 1.5 ml/min. A photo diode array was used as detector. Each component was identified by comparing its retention time and UV spectra with those of standard components. To further characterize the complexes, IR spectra and X-ray diffraction patterns were taken of them

Bulk polymerization

10

15

20

5

Propylene was polymerized in a stirred tank reactor having a volume of 5 l. About 0.9 ml triethyl aluminium (TEA) as a cocatalyst, about 0,12 ml of a 100-% solution of cyclohexyl methyl dimethoxy silane as an external donor and 30 ml of n-pentane were mixed and allowed to react for 5 minutes. Half of the mixture was added to the polymerization reactor and the other half was mixed with ca 20 mg of said catalyst component. After an additional 5 minutes the catalyst/TEA/donor/n-heptane mixture was introduced into the reactor. The Al/Ti mole ratio was 250 and the Al/external donor mol ratio was 10 mol/mol. 70 mmol of hydrogen and 1400 g of propylene were introduced into the reactor and the temperature was raised within 15-30 minutes to 70 °C. The polymerization time was 60 minutes, after which the polymer formed was taken out of the reactor. The polymers were characterized with respect to their Melt Flow Rate (MFR₂), bulk density (BD) and the fraction of total solubles in xylene (TS).

25 Results

Chemical composition of the catalysts

As stated in the experimental section, the catalysts were characterized with respect to their chemical composition. In Table 17 the chemical composition of the catalysts with respect to the Mg, Ti, di(2-ethyl-1-hexyl)phthalate (DOP), 2-ethyl-1-hexyl alcohol (EHA) and phthalic anhydride PA contents are listed in w-% units and in Table 18 the same species are listed in mol-% units and last, in Table 19 the molar composition between Mg, Ti and DOP are listed. The Examples 11 and 13 are represented by two catalysts, 11a and 11b, as well as 13a and 13b, respectively. The chlorine contents are listed in Table 20.

Table 18

Table 17

The Mg, Ti, DOP, EHA and PA contents of the catalysts in w-% units

Example	Mg w-%	Ti w-%	DOP w-%	EHA w-%	PA w-%
l la	3.6	5.6	35.6	6.8	4.16
11b	9.9	3.5	34.0	-	-
12	3.9	7.0	35.7	5.3	1.27
13a	4.5	7.0	43.6	5.15	1.6
13b	11.1	3.7	33.0	1.00	0.3

The Mg, Ti, DOP, EHA and PA contents of the catalysts in mol-% units

Example	Mg	Ti	DOP	EHA	PA
	mol-%	mol-%	mol-%	mol-%	mol-%
11a	0.148	0.117	0.091	0.052	0.028
11b	0.407	0.073	0.087	-	-
12	0.161	0.146	0.091	0.041	0.009
13a	0.185	0.146	0.112	0.040	0.011
13b	0.457	0.077	0.085	0.008	0.002

Table 19

Comparison between the molar amounts of Mg, Ti and DOP

Example	Mg/Ti	Ti	DOP/Ti
11a	1.3	1	0.8
11b	5.6	1	1.2
12	1.1	1	0.6
13a	1.3	1	0.8
13b	5.6	1	1.1

Table 20

The calculated amounts of Cl in the catalysts compared to the amounts found

Example	Calculated w-%	Found w-%
11a	27.1	26.0
11b	39	-
12	32.1	30.9
13a	33.5	32.4
13b	43.4	44.0

The IR results

In the IR spectra of the catalyst components of Examples 12 and 13a. There were clear indications of the presence of phthalic anhydride in the catalyst that has been prepared from MgR₂ (Example 11a). The phtalic anhydride was almost totally missing from the samples that had been prepared out of MgCl₂ (Examples 12 and 13a). These results confirm the results of the chemical analysis. The IR spectrum (not shown) for the toluene washed example 10b catalyst showed no traces of phthalic anhydride but to the left of the C=O---Ti peak a shoulder had appeared indicating the presens of some free carboxylic acid group (-COOH).

The X-ray diffraction patterns

The X-ray diffraction patterns for the catalysts show that the addition of TiCl₄ before PDC gives a more crystalline material. Example 12 is still showing the organic separation peak at 7° 2 Θ and the halo between 18° and 22° 2 Θ but only a slight remain thereof can be seen of the halo in the spectrum of Example 13a. In all patterns there seems to be an additional peak at about 32°-33° 2 Θ . This peak is not connected to crystalline MgCl₂. Some unreacted MgCl₂ seems to be present in the catalyst component of Example 13a which is starting to dominate when the catalyst is washed with toluene.

5

10

15

20

25

Polymerization results

All but one (Example 11a) of the catalysts were test polymerized according to the descriptions in the experimental section. The polymerization results both in kg PP/g cat units and in kg PP/g Ti units are listed in Table 21. There was an almost logaritmic linear increase in the activities. As a whole it can be said that:

 Addition of TiCl₄ before PDC gives better activity (compare Examples 12 and 13).

67

- 2. Starting from MgCl₂ instead of from MgR₂ gives higher activity (compare Example 11 with Examples 12 and 13).
 - 3. Toluene wash improves activity (compare Examples 13a and 13b).

The polymerization results

Example	Activity kg PP/g cat.	Activity kg PP/g Ti
11b	0.06	1.3
12	0.4	6.0
13a	1.2	18
134	1.2	10

2.5

Summary

13b

Table 21

15

In this study a stoichiometric synthesis route was used to produce the MgCl₂-C₆H₄(COOR')₂ complex. MgCl₂ and Mg-alkyl have been reacted with an alcohol to form a MgCl₂-Mg-alcoholate complex, a Mg-alcoholate and an MgCl₂-alcohol complex. These Mg-alcoholates or MgCl₂-alcohol complex has then been brought into contact with phthaloyl chloride to give an MgCl₂-donor complex. It was not possible to produce an MgCl₂-donor complex of this type through a direct contact between MgCl₂ and the corresponding donor, which rules out the possibility that this complex could have been formed unintactionally in the prior art. The complex achieved trough the synthesis routes described in this study is identifiable through its distinct X-ray diffraction pattern that shows a dominant peak at 4.5° 2Θ. IR studies showed that the Mg in the MgCl₂ is co-ordinated both to the C=O oxygen and the C-O- oxygen in the ester group.

Claims

5

10

15

1. A complex comprising a magnesium dihalide and an electron donor, characterized in that it is a complex of the magnesium dihalide and the electron donor and has the following formula (I) expressing the molar ratio between the magnesium dihalide and the electron donor:

$$MgX_2 \cdot [R(OR')_n]_m$$
 (I)

wherein MgX₂ is the magnesium dihalide and $R(OR')_n$ is the electron donor, X is a halogen, R is an n-valent C_1 - C_{20} aliphatic group, an n-valent C_7 - C_{27} araliphatic group or an n-valent C_2 - C_{22} acylic group, R' is a C_1 - C_{20} alkyl group or a C_7 - C_{27} aralkyl group, n is a number from 1 to 6 and m is defined as a number $0.5 \le m \le 2.0$.

- 2. The complex according to Claim 1, characterized in that X is selected from Cl, Br and I, and is preferably Cl.
- 3. The complex according to Claim 1 or 2, characterized in that R is an n-valent C₂-C₂₂ acylic group, preferably an n-valent aromatic C₇-C₂₂ acylic group, most preferably phthaloyl.
 - 4. The complex according to Claim 1, 2 or 3, characterized in that R' is a C₆-C₁₆ alkyl group, preferably a C₆-C₁₂ alkyl group like undecyl or 2-ethyl-1-hexyl.
- 5. The complex according to any preceding claim, characterized in that n is 1 to 20 4, preferably about 2.0.
 - 6. The complex according to any preceding claim, characterized in that m is 0.67 to 1.0.
 - 7. The complex according to any preceding claim, characterized in that it is a magnesium dichloride phthalic acid ester complex having the formula (II):

25
$$MgCl_2 \cdot C_6H_4(COOR')_2$$
 (II)

wherein R' is the same as above.

8. The complex according to one of Claims 1 to 5, characterized in that it is a magnesium dichloride phthalic acid ester complex having the formula (III):

$$(MgCl2)3 \cdot [C6H4(COOR')2]2$$
 (III)

10

wherein R' is the same as above.

- 9. The complex according to any preceding claim, characterized in that it has an X-ray diffraction pattern showing a dominant peak at 4.5°2Θ.
- 10. Process for the preparation of a complex according to any preceding claim comprising a magnesium dihalide and an electron donor, characterized by reacting a magnesium compound (a) containing an alkoxy moiety, which magnesium compound is selected from the group consisting of a complex of a magnesium dihalide and a magnesium dialkoxide, a complex of a magnesium dihalide and an alcohol, and a non-complex magnesium dialkoxide, with a halogen compound (b), which is capable of forming the electron donor by replacement of its halogen by said alkoxy moiety.
 - 11. Process according to Claim 10, characterized in that said halogen compound (b) has the formula (IV):

RXn (IV)

- wherein R is an n-valent C₁-C₂₀ aliphatic group, an n-valent C₇-C₂₇ araliphatic group or an n-valent C₂-C₂₂ acylic group, X is a halogen and n is 1 to 6.
 - 12. The complex according to Claim 11, characterized in that R is an n-valent C_2 - C_{22} acylic group, preferably an n-valent aromatic C_7 - C_{22} acylic group, most preferably phthaloyl.
- 20 13. Process according to Claim 11 or 12, characterized in that X is selected from Cl, Br and I, and is preferably Cl.
 - 14. Process according to Claim 11, 12 or 13, characterized in that n is 1 to 4, preferably about 2.
- 15. Process according to any of Claims 11 to 14, characterized in that said halogen compound is an organic acid halide, preferably phthalic acid dichloride Ph(COCl)2, wherein Ph is o-phenylene.
 - 16. Process according to any of Claims 10 to 15, characterized in that said complex of a magnesium dihalide and a magnesium dialkoxide is a magnesium dichloride-magnesium dialkoxide complex of the formula (V):

 $MgCl_2 \cdot [Mg(OR')_2]_p$

wherein R' is a C_1 - C_{20} alkyl group or a C_7 - C_{27} aralkyl group, preferably a C_6 - C_{16} alkyl group, and p is 1 to 6, preferably about 2.

17. Process according to Claim 16, characterized in that said complex of a magnesium dihalide and a magnesium dialkoxide is a magnesium dichloride-dimagnesium dialkoxide complex of the formula (VI):

$$MgCl_2 \cdot [Mg(OR')_2]_2$$
 (VI)

wherein R' is a C_1 - C_{20} alkyl group or a C_7 - C_{27} aralkyl group, preferably a C_6 - C_{16} alkyl group.

- 18. Process according to Claim 16, characterized in that said magnesium dichloride magnesium dialkoxide complex is prepared by reacting magnesium dichloride with and alcohol into an intermediate which is a magnesium dichloride alcohol complex MgCl₂·(R'OH)_{2p}, wherein R' is the same as above, and reacting the magnesium dichloride alcohol complex with p mol of a magnesium dialkyl MgR"₂, wherein R" is a hydrocarbyl group having 1 to 20 carbon atoms.
- 15 19. Process according to Claim 18, characterized in that, independently, the molar ratio MgCl₂:R'OH is between 1:1 and 1:8, preferably between 1:2 and 1:5, the molar ratio MgCl₂·(R'OH)_{2p}:MgR"₂ is between 1:1 and 1:4, preferably about 1:2, the temperature is between 80 °C and 160 °C, and the reaction time is about 2 h to about 8 h.
- 20. Process according to Claim 15 and 17, characterized in that said magnesium compound (a) which is said magnesium dichloride-dimagnesium dialkoxide complex MgCl₂·[Mg(OR')₂]₂, wherein R' is a C₆-C₁₆ alkyl group, is reacted with said halogen compound (b) which is said phthalic acid dichloride Ph(COCl)₂, wherein Ph is o-phenylene.
- 25 21. Process according to any of Claims 10 to 15, characterized in that said non-complex magnesium dialkoxide has the formula (VII):

$$Mg(OR')_2$$
 (VII)

wherein R' is a C₁-C₂₀ alkyl group or a C₇-C₂₇ aralkyl group, preferably a C₆-C₁₆ alkyl group.

30 22. Process according to Claim 21, characterized in that said non-complex magnesium dialkoxide is prepared by reacting a magnesium dialkyl, preferably a

20

25

5

magnesium dialkyl of the formula MgR"2, wherein R" is a hydrocarbyl group having 1 to 20 carbon atoms, and an alcohol, preferably an alcohol of the formula R'OH wherein R' is the same as above.

- 23. Process according to Claim 15 and 21, characterized in that said magnesium compound (a) which is said non-complex magnesium dialkoxide has the formula Mg(OR')2, wherein R' is a C₁-C₂₀ alkyl or a C₇-C₂₇ aralkyl, preferably a C₆-C₁₆ alkyl, is reacted with said halogen compound (b) which is said phthalic acid dichloride Ph(COCl)2, wherein Ph is o-phenylene.
- 24. Process according to any of Claims 10 to 15, characterized in that said complex of a magnesium dihalide and a magnesium dialkoxide is a complex of a magnesium dichloride and an alcohol having the formula (VIII):

$$MgCl_2 \cdot (R'OH)_q$$
 (VIII)

wherein R' is a C_1 - C_{20} alkyl group or a C_7 - C_{27} aralkyl group, preferably a C_6 - C_{16} alkyl group, and q is from 1 to 6.

- 25. Process according to Claim 24, characterized in that said complex of a magnesium dihalide and an alcohol is prepared by reacting magnesium dichloride MgCl₂ and and alcohol R'OH, wherein R' is the same as above.
 - 26. Process according to Claim 24 or 25, characterized in that the reaction temperature is kept between 10 °C and 100 °C, and the reaction time is about from 10 to about 90 min.
 - 27. Process according to Claim 15 and 24, characterized in that said magnesium compound (a) which is said complex of a magnesium dihalide and an alcohol having the formula $MgCl_2 \cdot (R'OH)_q$, wherein R' is a C_1 - C_{20} alkyl or a C_7 - C_{27} aralkyl, preferably a C_6 - C_{16} alkyl, and q is from 1 to 6, is reacted with said halogen compound (b) which is said phthalic acid dichloride $Ph(COCl)_2$, wherein Ph is ophenylene.
 - 28. Process according to one of Claims 10 to 27, characterized in that said magnesium compound (a) and said halogen compound (b) are reacted essentially stoichiometrically.
- 30 29. Use of a complex according to one of Claims 1 to 9 or a complex prepared according to one of Claims 10 to 28 for the preparation of a polymerization catalyst component containing magnesium, transition metal, halogen and electron donor.

- 30. Use according to Claim 29, characterized in that said complex is reacted with a titanium halide (c).
- 31. Use according to Claim 30, characterized in that said titanium halide (c) has the formula (IX):

$$5 \qquad (OR''')_{p}TiX_{4-p} \qquad (IX)$$

wherein R'' is a C_1 - C_{10} alkyl group or a C_7 - C_{16} aralkyl group, X is a halogen and p is 0 to 3, and preferably is a titanium tetrahalide TiX₄, wherein X is the same as above, most preferably titanium tetrachloride TiCl₄.

32. A complex according to one of claims 1 to 9, characterized in that it shows an IR spectra with a main absorption peak at 1729 cm⁻¹ for the C=O...Mg that has shifted 5 to 15 cm⁻¹, preferably 10 cm⁻¹ to the right, and preferably also shows three shoulders.

(57) Abstract

The invention relates to a complex comprising a magnesium dihalide and an electron donor. It is a complex of the magnesium dihalide and the electron donor and has the formula (I):

$$MgX_2 \cdot [R(OR')_n]_m$$
 (I)

wherein MgX₂ is the magnesium dihalide and R(OR')_n is the electron donor, X is a halogen, R is an n-valent C_1 - C_{20} aliphatic group, an n-valent C_7 - C_{27} araliphatic group or an n-valent C_2 - C_{22} acylic group, R' is a C_1 - C_{20} alkyl group or a C_7 - C_{27} aralkyl group, n is a number from 1 to 6 and m is defined as a number $0.5 \le m \le 2.0$. The invention also relates to the preparation process of such a complex, as well as the use thereof for the preparation of olefin polymerization catalyst components.

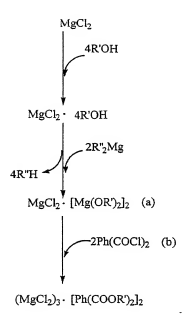


Figure 1 Example of first alternative embodiment

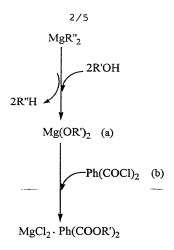


Figure 2 Example of second alternative embodiment

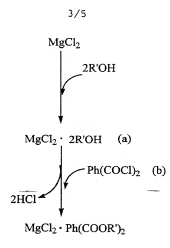


Figure 3 Example of third alternative embodiment

الم المنظم ا

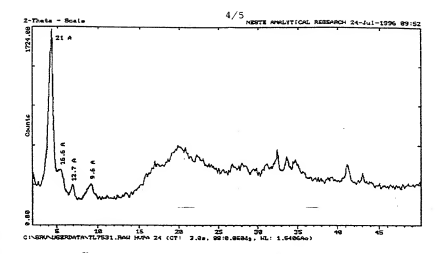


Figure 4 The x-ray diffraction pattern of the Mg-complex produced out of Mg-alkyl, alcohol and phthalic ester (example 2)

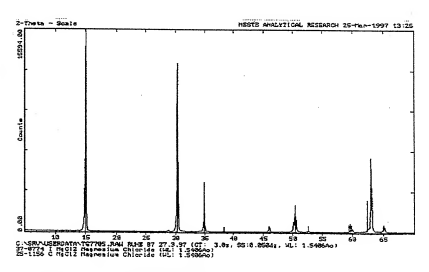


Figure 5 The x-ray diffraction pattern of the product of MgCl₂ and phthalic ester (comparative example)

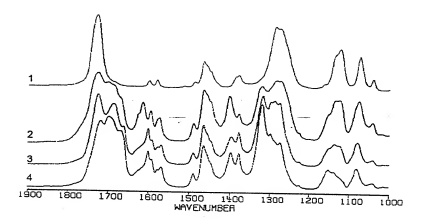


Figure 6 IR Spectra of

1 DÛP,

2 MgCl₂ · DOP (example 2),

3 (MgCl₂)_{1.5} · DUP (example 1) and

4 (MgCl₂)_{1.5} · DUP (example 1) stored overnight

F السؤ 101

PLEASE NOTE: YOU MUST COMPLETE THE

COMBINED DECLARATION AND POWER OF ATTORNEY FOR PATENT APPLICATION

ATTORNEY DOCKET NO.

696-171PCT

As a below named inventor, I hereby declare that: my residence post office address and citizenship are as stated next to my name; that I verily believe that I am the original, first and sole inventor (if only one inventor is named below) or a joint inventor (if plural inventors are named below) of the subject matter which is claimed and for which a patent is sought on the invention entitled: *

	Soluble magnesium dihalide complex, prepar	ation and	use
Box II			IPE
e Without	the specification of which is attached hereto unless one of the follow	ing boxes below	් පි checked ්ර
ration 🖈	The Specification was filed on	and was as∮igr	ned anna
	Serial No.09/582321 and was amended on		AUG 2 5 2000 on
	was filed as PCT international application number		on &
	and was amended under PCT Article 19 on		by city
	(if applicable).		TRADEMAR

I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, §1.56(a).

I do not know and do not believe the same was ever known or used in the United States of America before my or our invention thereof, or patented or described in any printed publication in any country before my or our invention thereof, or more than one year prior to this application, that the same was not in public use or on sale in the United States of America more than one year prior to this application, that the invention has not been patented or made the subject of an inventor's certificate issued before the date of this application in any country foreign to the United States of America on an application filed by me or my legal representatives or assigns more than twelve months prior to this application, and that no application for patent or inventor's certificate on this invention has been filed in any country foreign to the United States of America prior to this application by me or my legal representatives or assigns, except as follows:

I hereby claim foreign priority benefits under Title 35, United States Code, §119 of any foreign application(s) for patent or inventor's certificate listed below:

Prior Foreign Application(s)			Priority	Claimed
974622 (Number)	Finland (Country)	12/23/1997 (Month/ Day: Year Filed)	X Yes	□ No
974623 (Number)	Finland (Country)	12/23/1997 (Month, Day Year Filed)	X⊟ Yes	□ No
(Number)	(Country)	(Month/Day, Year Filed)	□ Yes	□ No
(Number)	(Country)	(Month/Day Year Filed)	Yes	□ No
(Number)	(Country)	(Month/ Day: Year Filed)	Yes	□ No

Months Prior To The Filing Date of This Application:

Country	Application No.	Date of Filing (Month/Day: Year)
PCT	PCT/FI98/01004	12/21/1998

I hereby claim the benefit under Title 35, United States Code, §120 of any United States application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35, United States Code, §112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, §1.56(a) which occurred between the filing date of the prior application and the national or PCT international filing date of this application:

(Application Serial No.)	(Filing Date)	(Status - patented, pending, abandoned)
(Application Serial No.)	(Filing Date)	(Status — patented, pending, abandoned)

I hereby appoint the following attorneys to prosecute this application and/or an international application based on this application and to transact all business in the Patent and Trademark Office connected therewith and in connection with the resulting patent based on instructions received from the entity who first sent the application papers to the attorneys identified below, unless the inventor(s) or assignee provides said attorneys with a written notice to the contrary:

 RAYMOND C. STEWART (Reg. No. 21.066)
 TERRELL C. E

 JOSEPH A. KOLASCH (Reg. No. 22.463)
 ANTHONY L.

 JAMES M. SLATTERY (Reg. No. 28.380)
 BERNARD L.

 DONALD C. KOLASCH (Reg. No. 23.038)
 MICHAEL K. I

 CHARLES GORENSTEIN (Reg. No. 29.071)
 GERALD M. N

 LEONARD R. SVENSSON (Reg. No. 30, 330)

TERRELL C. BIRCH (Reg. No. 19.382)
ANTHONY L. BIRCH (Reg. No. 26,122)
BERNARD L. SWEENEY (Reg. No. 24,448)
MICHAEL K. MUTTER (Reg. No. 29,680)
GERALD M. MURPHY, JR. (Reg. No. 28,977)

Send Correspondence to: BIRCH, STEWART, KOLASCH AND BIRCH

301 North Washington Street P.O. Box 747

Falls Church, Virginia 22040-0747 Telephone: (703) 241-1300

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

DATE GIVEN NAME FAMILY NAME INVENTOR'S SIGNATURE 20 Mcs Raro H 10 Aug. 2000 GAROFF Thomas RESIDENCE (City, State & Country) Helsinki Finland Finland POST OFFICE ADDRESS (Complete Street Address including City, State & Country) Kirkkosalmentie 6 A 5, FIN-00840 Helsinki GIVEN NAME FAMILY NAME INVENTOR'S SIGNATURE DATE Timo LEINONEN 10 Aug. 2000 RESIDENCE (City, State & Country) CITIZENSHIP Tolkkinen Finland Finland POST OFFICE ADDRESS (Complete Street Address Massatie 8, FIN-06750 Tolkkinen.Finland GIVEN NAME FAMILY NAME INVENTOR'S SIGNATURE DATE Sirpa ALA-HUIKKU 10 Aug. 2000 RESIDENCE (City, State & Country) CITIZENSHIP Helsinki, Finland Finland POST OFFICE ADDRESS (Complete Street Address including City, State & Country) Korvatunturintie 5 B, FIN-00970 Helsinki. INVENTOR'S SIGNATURE GIVEN NAME FAMILY NAME DATE RESIDENCE (City, State & Country) CITIZENSHIP POST OFFICE ADDRESS (Complete Street Address including City, State & Country) GIVEN NAME FAMILY NAME INVENTOR'S SIGNATURE *DATE CITIZENSHIP RESIDENCE (City, State & Country) POST OFFICE ADDRESS (Complete Street Address including City, State & Country)

3-00